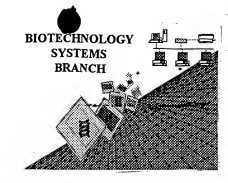
RAW SEQUENCE LISTING ERROR REPORT



The Biotechnology Systems Branch of the Scientific and Technical Information Center (STIC) detected errors when processing the following computer readable form:

Application Serial Number:	09/676,/35
Source:	OIPE
Date Processed by STIC:	10/11/2000

THE ATTACHED PRINTOUT EXPLAINS DETECTED ERRORS.
PLEASE FORWARD THIS INFORMATION TO THE APPLICANT BY EITHER:

- 1) INCLUDING A COPY OF THIS PRINTOUT IN YOUR NEXT COMMUNICATION TO THE APPLICANT, WITH A NOTICE TO COMPLY or,
- 2) TELEPHONING APPLICANT AND FAXING A COPY OF THIS PRINTOUT, WITH A NOTICE TO COMPLY

FOR CRF SUBMISSION QUESTIONS, PLEASE CONTACT MARK SPENCER, 703-308-4212.

FOR SEQUENCE RULES INTERPRETATION, PLEASE CONTACT ROBERT WAX, 703-308-4216. PATENTIN 2.1 c-mail help: patin21help@uspto.gov or phone 703-306-4119 (R. Wax) PATENTIN 3.0 c-mail help: patin30help@uspto.gov or phone 703-306-4119 (R. Wax)

TO REDUCE ERRORED SEQUENCE LISTINGS, PLEASE USE THE CHECKER VERSION 3.0 PROGRAM, ACCESSIBLE THROUGH THE U.S. PATENT AND TRADEMARK OFFICE WEBSITE. SEE BELOW:

Checker Version 3.0

The Checker Version 3.0 application is a state-of the-art Windows based software program employing a logical and intuitive user-interface to check whether a sequence listing is in compliance with format and content rules. Checker Version 3.0 works for sequence listings generated for the original version of 37 CFR §§1.821 – 1.825 effective October 1, 1990 (old rules) and the revised version (new rules) effective July 1, 1998 as well as World Intellectual Property Organization (WIPO) Standard ST.25.

Checker Version 3.0 replaces the previous DOS-based version of Checker, and is Y2K-compliant. Checker allows public users to check sequence listings in Computer Readable form (CRF) before submitting them to the United States Patent and Trademark Office (USPTO). Use of Checker prior to filing the sequence listing is expected to result in fewer errored sequence listings, thus saving time and money.

Checker Version 3.0 can be down loaded from the USPTO website at the following address: http://www.uspto.gov/web/offices/pac/checker

OIPE

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10/11/00

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DATE: 10/11/2000
                   RAW SEQUENCE LISTING
                   PATENT APPLICATION: US/09/676,135
                                                        TIME: 12:29:07
                   Input Set : A:\ES.txt
                   Output Set: N:\CRF3\10112000\I676135.raw
                                                                                  Does Not Comply
                                                                            Corrected Diskette Needed
     3 <110> APPLICANT: Boyle, Brian J
            Mize, Nancy K
            Arterburn, Matthew C
     6
            Yeung, George
            Tang, Y. Tom
     8
            Liu, Chenghua
            Drmanac, Radoie T
            Wang, Menq-Yun
    10
    11
            Chen, Lichuan
    12
            Yang, Yea-Huey
    14 <120> TITLE OF INVENTION: METHODS AND MATERIALS RELATING TO METALLOCARBOXYPEPTIDASE-LIKE
    15
            POLYPEPTIDES AND POLYNUCLEOTIDES
    17 <130> FILE REFERENCE: HYS-28
    19 <140> CURRENT APPLICATION NUMBER: US/09/676,135
    20 <141> CURRENT FILING DATE: 2000-09-29
    22 <150> PRIOR APPLICATION NUMBER: US 09/560,875
    23 <151> PRIOR FILING DATE: 2000-04-27
    25 <150> PRIOR APPLICATION NUMBER: US 09/496,914
    26 <151> PRIOR FILING DATE: 2000-02-03
    28 <160> NUMBER OF SEQ ID NOS: 22
    30 <170> SOFTWARE: PatentIn version 3.0
ERRORED SEQUENCES
                                                 why is a gap present? Per 1.822(e)
at the Tyr Glu The Sequence Rules,
    565 <210> SEQ ID NO. 19 /64 shows
    567 <212> TYPE: PRT
    568 <213> ORGANISM: homo sapiens
    570 <400> SEQUENCE: 19
                                             Val Thr Tyr Glu Thr
    572 Tyr Lys Glu Val Val Thr Gln His Phe Leu
E--> 573 1
                                         10
    575 His Pro Ile Tyr Tyr Leu Lys Ile Ser Gln Pro Ser Gly Asn Pro Lys
                                                                         "A sequene with a gap
                   20
E--> 576
    578 Lys Ile Ile Trp Met Asp Cys Gly Ile His Ala Arg Glu Trp Ile Ala
E--> 579
              3.5
                                 40
                                                   45
                                                                         or gaps shall be
    581 Pro Ala Phe Cys Gln Trp Phe Val Lys Glu Ile Leu Gln Asn His Lys
                              55
                                                60
  -> 582
    584 Asp Asn Ser Arg Ile Arg Lys Leu Leu Met Asn Leu Asp Phe Tyr Val
                                                                             esented as a plurality
E--> 585 65
                          70
    587 Leu Pro Val Leu Asn Ile Asp Gly Tyr Ile Tyr Thr Trp Thr Thr Asp
E--> 588
                       8.5
                                         90
    590 Arg Leu Trp Arg Lys Ser Arg Ser Pro His Asn Asn Gly Thr Cys Phe
                                    105
                                                        110
E--> 591
    593 Gly Thr Asp Leu Asn Arg Asn Phe Asn Ala Ser Trp Cys Ser Ile Gly
                                                                        with separate sequence
                                 120
E--> 594
               115
    596 Ala Ser Arg Asn Cys Gln Asp Gln Thr Phe Cys Gly Thr Gly Pro Val
                                                                      rdestifier, with the
                                                                     number of separate
                                                                       sequerar being equal in
                                                                     number to the number of
                                                                      continuous strings of
                                                                     sequence deta, el gap
                                                                   represents a stop codon,
                                                                    arend ande following
```

RAW SEQUENCE LISTING

DATE: 10/11/2000 TIME: 12:29:07

PATENT APPLICATION: US/09/676,135

Input Set : A:\ES.txt

Output Set: N:\CRF3\10112000\1676135.raw

> 597 130 135 140 599 Ser Glu Pro Glu Thr Lys Ala Val Ala Ser Phe Ile Glu Ser Lys Asn 150 155 160 E--> 597 130

E--> 600 145 150

602 Asp Asp Phe Cys Ala 165 E--> 603

VERIFICATION SUMMARY PATENT APPLICATION: US/09/676,135

DATE: 10/11/2000 TIME: 12:29:08

Input Set : A:\ES.txt

Output Set: N:\CRF3\10112000\1676135.raw

L:19 M:270 C: Current Application Number differs, Replaced Current Application Number L:573 M:332 E: (32) Invalid/Missing Amino Acid Numbering, SEQ ID:19 M:332 Repeated in SeqNo=19 L:603 M:252 E: No. of Seq. differs, <211>LENGTH:Input:165 Found:164 SEQ:19 L:748 M:257 W: Feature value mis-spelled or invalid, <221> Name/Key for SEQ ID#:22 L:755 M:341 W: (46) "n" or "Xaa" used, for SEQ ID#:22